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gp100 gene. This fragment was digested with *EcoRV* and *Xba*I and cloned into *EcoRV/Xba*I digested NVQH6MC5#10 generating plasmid C5H6MELgp100 #5 which contains the gp100 gene linked to the H6 promoter.

The gp100 gene in plasmid C5H6MELgp100 #5 was sequenced using 5 custom primers. A 65bp deletion was found in this clone and shown to be present in pCDNA3-gp100. Plasmid PCRII-gp100 was used in PCR with oligonucleotides MELgp05(5'-CCC-ATC-TGG-CTC-TTG-GTC-3') (SEQ.ID.NO. 115) and MELgp13 (5'-TGA-CAT-CTC-TGC-CAG-TGT-GGT-3') (SEQ.ID.NO. 116) to generate a 0.6kb fragment. This fragment was digested with *Bam*HI and 10 *Asp*718 and ligated to a 6.5kb *Asp*718/*Bam*HI (partial) fragment from C5H6MELgp100 #5 generating plasmid C5H6MELgp100 which contains the entire gp100 gene under the control of the H6 promoter.

Pre-existing plasmid pC5H6MELgp100 was used as template for site directed mutagenesis of the two CTL epitopes beginning at amino acids 209 and 15 280, respectively. Primers used were:

209-A

GCT CAG CCT TCA CCA TTA TGG ACC AGG TGC CTT TCT CC
(SEQ.ID.NO.117)

209-B

20 GGA GAA AGG CAC CTG GTC CAT AAT GGT GAA GGC TGA CG
(SEQ.ID.NO.118)

280-A

GAG CCT GGC CCA GTC ACT GTT CAG GTG GTC CTG CAG GC
(SEQ.ID.NO.119)

25 280-B

GCC TGC AGG ACC ACC TGA ACA GTG ACT GGG CCA GGC TC
(SEQ.ID.NO.120)

A section containing the modified epitopes was sequenced and isolated as a 440 bp *Nco*I/*Mlu*I fragment. This fragment was ligated into

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pC5H6MELgp100 digested with *Nco*I and *Mlu*I, creating a plasmid with the complete gp100 with the modified epitopes 209-2M and 280-9V.

Sequence data revealed a G to C substitution at bp# 10, changing a.a. # 4 from a Valine to a Leucine. This was corrected by PCR using the following

5 primer pair:

MEL25

GCT CCG GGA TCC CCG GCC ATG GTA GAC AGT CAC TTC CAT CGT GTG
TGT GCC CAG CAT TG (SEQ.ID.NO.121)

MEL27

10 ATC GCG ATA TCC GTT AAG TTT GTA TCG TAA TGG ATC TGG TGC TAA
AAA GAT GCC TTC TT (SEQ.ID.NO.122)

MEL25 changes bp# 549 from a C to a G destroying the unique *Nco*I site for easier screening. It does not change the amino acid.

The resulting PCR fragment was digested with *Bam*H1 and *Eco*R5 and
15 replaced the equivalent fragment correcting the error. The resulting plasmid is pC5gp100-M which is shown in Figure 3 (SEQ.ID.NO.123).

Genetic modification of the recipient:

Recombination between donor plasmid pC5gp100M and ALVAC(2) rescuing virus generated recombinant virus vCP1584, which contains the
20 vaccinia H6 promoted modified human gp100 in the C5 locus.

EXAMPLE 3

Screening for the identification and purification of recombinant organisms:

The aspects of screening for the identification and purification of a recombinant organism of the present invention is set out below.

25 (1) Plaque purification was done using *in situ* plaque hybridization (Piccini *et al.*, Methods of Enzymol. 153:545 (1987)) was used to identify recombinant viruses and to demonstrate purity of final virus preparations. *In situ* plaque hybridization analysis was performed with radiolabelled probes specific for the gp100 construct (a 580 bp fragment) and the C5 insertion locus.

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(2) Restriction analysis: Viral genomic DNA was isolated from cells infected with ALVAC parent or ALVAC(2)-gp100M (vCP1584). The genomic DNA was digested with restriction endonucleases (*Hind*III, *Pst*I or *Bam*HI). The resultant DNA fragments were fractionated by electrophoresis through an agarose gel and visualized by ethidium bromide staining. The insertion of the mod gp100 expression cassette at the C5 locus was confirmed.

(3) Immunoprecipitation analyses: These were performed using radiolabeled lysates derived from uninfected HeLa cells or cells infected with either ALVAC parental virus, ALVAC-gp100 (vCP1465) or ALVAC(2)-gp100M (vCP1584) as described previously (Taylor *et al.* J. Virol. 64:1441 (1990)). Briefly, HeLa cell cultures were infected at an m.o.i. of 10 pfu/cell in methionine-free media supplemented with [35S]-methionine (35uCi/ml). At 18 hrs. post infection, cells were lysed. Immunoprecipitation was performed using a rabbit anti-gp100 serum (AZN-LAM, received from M. Schreurs University of Nijmegen, Netherlands). Immunoprecipitates were fractionated on a 10% SDS-Polyacrylamide gel. The gel was fixed and treated for fluorography with 1M Na-salicylate for 1/2 hr. The dried gel was exposed to Kodak XAR-2 film to visualize the protein species. Results with anti-gp100 demonstrate expression of gp100 in ALVAC-gp100 infected HeLa cells but not for parentally infected cells. (See Figure 6)

(4) Western Blot: HeLa cells were infected for 18 hours at a multiplicity of 10 pfu/cell with ALVAC(2)-gp100M (vCP1584), ALVAC-gp100 (vCP1465) or ALVAC. Cell lysates were separated by SDS-PAGE and transferred to nitrocellulose. The blot was incubated with AZN-LAM (1/5000 dilution) followed by HRP conjugated swine anti-rabbit utilizing the enhanced chemiluminescence (ECL) detection method (Amersham). Results demonstrate expression of full length gp100 in ALVAC-gp100 and ALVAC(2)-gp100M infected cells. (See Figure 7).

(5) Plaque immunoscreen analysis: This was performed on vCP1584 material to determine phenotypic stability of the virus upon passaging. The

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phenotypic stability of production batch material of ALVAC-gp100M (vCP1584) was analyzed by an immunological plaque assay which measures expression of the inserted genes at the plaque level. The assay utilizing permeabilized cells for detection of intracellular as well as surface expression of Hgp100mod was chosen
5 for this test.

Test and control reagents (ALVAC(2)-gp100M (vCP1584) and ALVAC standard and ALVAC-gp100M, respectively) were plated on CEF monolayers under agarose at dilutions resulting in 40-200 plaques per 60 mm dish. 120 hours after incubation at 37°C, the infected monolayers were processed by plaque
10 immunoassay for detection of internal expression of gp100M. Positive and negative plaques were counted for test and control samples. The primary antibody used was Monoclonal Anti-HMB50 at 1:800 dilution. A secondary antibody used was horse radish peroxidase (HRP)-conjugated rabbit anti-mouse antiserum diluted 1:500.

15 The result of analysis of internal expression of Human modified gp100 by individual plaques produced by (vCP1584) is presented in Table 1.

The result demonstrates that 98.7% of the plaque population of ALVAC-gp100M is expressing gp100M indicating that ALVAC-gp100M is phenotypically stable.

20 Results of the plaque immunoscreen analysis demonstrate that ALVAC(2)-gp100M is phenotypically stable with respect to expression of gp100.

(6) Nucleotide sequence analysis. This was performed on vCP1584 to validate the nucleotide sequence of the H6-promoted melanoma gp100M cassette. The sequence analysis revealed no nucleotide differences relative to the
25 expected sequence, thus no mutations were introduced during the production of vCP1584. In order to carry out this analysis, a pool of plasmid clones containing a 2.2 kb PCR-derived fragment (encompassing the H6-promoted melanoma gp100M cassette), generated from vCP1584 genomic DNA was used.

pBS/1584 was generated by pooling 9 positive clones obtained by the
30 ligation of a 2.2 kb PCR fragment (containing the H6-promoted melanoma

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gp100M cassette from vCP1584), into pBS-sk-(Stratogene). The 2.2 kb PCR fragment was derived from vCP1584 genomic DNA with the oligonucleotide primers, IDC5-1 and IDC5-2 (Figure 5). The nucleotide sequence of the oligonucleotide primers used to sequence pBS/1584 are listed in Figure 5.

5 **EXAMPLE 4**

This example provides results from injection in cynomolgus monkeys of modified gp100 molecules.

Methods and Experimental Design

Test System

- 10 Cynomolgus monkeys (*Macaca fascicularis*) purpose bred animals.

Supplier: Siconbrec "Simian Conservation Breeding & Research Center Inc.", Fema Building, 44 Gil Puyat Avenue Makati, Metro Manila, Philippines.

Number of animals in the study: 12 (6 males and 6 females).

Age at initiation of treatment: 26 to 38 months.

- 15 - Body weight range at initiation of treatment (day -1):

- males: 1.73 to 2.34 kg

- females: 1.71 to 2.65 kg.

Animal Husbandry

- Housing: one air-conditioned room;

- 20 - temperature: 19 to 25°C (target range),

- relative humidity: >40%

- air changes: minimum 8 air changes per hour,

- lighting cycle: 12 hours light (artificial)/12 hours dark.

- 25 - Caging: animals were housed singly in stainless steel mesh cages (approximately 540 x 810 x 760 mm).

- Diet: expanded complete commercial primate diet (Mazuri diet, Special Diet Services Ltd., Witham, Essex, CM8, 3AD, Great Britain) analyzed for chemical and bacterial contaminants.

Quantity distributed: 100g diet/animal/day.

- 30 In addition, animals received fruit daily (apple or banana)

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Animals were fasted for at least 16 hours before blood sampling for clinical laboratory investigations and before necropsy.

- Water: drinking water *ad libitum* (via bottles).
- Contaminants: no known contaminants were present in diet or water at levels which might have interfered with achieving the objective of the study.

Pre-Treatment Procedures

- Animal health procedure: all animals received a clinical examination for ill-health on arrival and a veterinary clinical examination during the acclimatization period.
- 10 - Acclimatization period: at least 3 weeks between animal arrival and start of treatment.

Experimental Design

- Allocation to treatment groups was performed during the acclimatization period using a random allocation procedure based on body weight classes.
- 15 - Animals were assigned to the treatment groups shown in Table 2. The dose levels administered were shown in Table 3.

Administration of the Test/Control Articles

Group 1 and 2 Animals

- Method of administration: injection in the left inguinal lymph node. Animals were lightly anaesthetized before each administration by an intramuscular injection of ketamine hydrochloride (Imalgene® 500 - Merial, Lyon, France). The same lymph node was injected on each occasion (left side). Each injection was followed by a local disinfection with iodine (Vétédine® - Vétoquinol, Lure, France).

Group 3

- Route: subcutaneous.
- Method of administration: bolus injection using a sterile syringe and needle introduced subcutaneously. Four injection sites were used followed by a local disinfection with iodine (Vétédine® - Vétoquinol, Lure, France). Animals were also lightly anaesthetized before each administration by an intramuscular

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injection of ketamine hydrochloride (Imalgene® 500 - Merial, Lyon, France) in order to be under the same conditions as groups 1 and 2 animals.

Four injection sites in the dorsal cervical/interscapular regions were used as shown in Table 4.

5 **ELISPOT Analysis**

An ELISPOT assay was used in order to assess the cell mediated immune response generated in the monkeys in the various treatment groups. In particular, an ELISPOT IFN γ assay was used in order to measure IFN γ production from T lymphocytes obtained from the monkeys in response to

10 gp100 antigens.

Materials and Methods

Plates: MILLIPORE Multiscreen HA plate / MAHA S45.10 (96 wells).

Capture antibodies: MABTECH monoclonal anti-IFN γ antibodies/G-Z4 1 mg/mL.

15 Detection antibodies: MABTECH monoclonal anti-IFN γ antibodies/7-B6-1-biotin 1 mg/mL.

Enzyme: SIGMA, Extravidin-PA conjugate/E2636

Substrate: BIORAD, NBT/BCIP - Alkaline phosphatase conjugate substrate kit/ref: 170-64 32.

20 **Coating**

Place 100 μ L per well of capture antibodies at 1 μ g/mL diluted at 1/1000 in carbonate bicarbonate buffer 0.1M pH 9.6 into the multiwell plate. Incubate overnight at 4°C. Wash 4 times in 1X PBS.

Saturation

25 Place 200 μ L per well of RPMI supplemented with 10% FCS, non essential amino acids, pyruvate, Hepes buffer and Peni-Strepto. Incubate 2 hours at 37°C.

Test

Cells from the immunized animals are tested against (a) medium alone; (b) pooled peptides at a concentration of 1 mg/mL; and (c) a non specific stimulus

30 (PMA-Iono). The pooled peptides used in this Example to stimulate IFN- γ

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production were derived from gp100 and are illustrated in Tables 5 to 8. The final volume of each sample is 200 µL. Incubate 20 hours at 37°C.

Wash 4 times in 1X PBS and 0.05% Tween 20.

Detection

- 5 Place 100 µL per well of detection antibodies at 1 µg/mL diluted in 1/1000 1X PBS, 1% BSA and 0.05% Tween 20. Incubate 2 hours at room temperature. Wash 4 times in 1X PBS and 0.05% Tween 20.

Reaction

- Place 100 µL per well of Extravidin-PA conjugate diluted 1/6000 in 1X PBS, 1%
10 BSA and 0.05% Tween 20. Incubate 45 minutes at room temperature. Wash 4 times in 1X PBS and 0.05% Tween 20.

Substrate Addition

- Place 100 µL per well of substrate previously prepared. For example, for 1 plate, prepare: 9.6 mL of distilled water, 0.4 mL of 25X buffer, 0.1 mL of solution A
15 (NBT) and 0.1 mL of solution B (BCIP). Incubate 30-45 minutes at room temperature. Wash in distilled water. Dry and transfer to a plastic film. The number of spots are counted using a Zeiss image analyzer. Each spot corresponds to an individual IFN-γ secreting T cell.

Results

- 20 The results of the ELISPOT analysis are shown in Figures 8-11. The results demonstrate that of the animals tested, 2 out of 2 (i.e. 100%) of the animals that received the intranodal administration of the gp100 antigen, and 2 out of 4 (i.e. 50%) of the animals that received the subcutaneous administration of the gp100 antigen had a positive cell mediated immune response.

25 ELISA Analysis

- The ELISA was performed utilizing standard methodology known in the art. Briefly, the human gp100 ("hgp100"; produced in Baculovirus) was diluted in coating buffer (carbonate-bicarbonate, pH9.6) and added to 96 wells at 0.5µg/well. Plates were placed at 4°C overnight. Plates were then washed and 30 blocking buffer (phosphate buffered saline/0.5% Tween 20/1.0% BSA, pH7.2)

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was added for 2 hours at 37°C. The plates were then washed and the sera was diluted in dilution buffer (phosphate buffered saline/0.5 % Tween 20/ 0.1 BSA, pH7.2). For this study, monkey sera was diluted to 1:800 and "7" serial 3 fold dilutions were done for each sample tested. The human sera controls were
5 diluted to 1:50 in dilution buffer and "7" serial 2 fold dilutions were performed. Each dilution was done in duplicate. The plates were incubated a further 2 hours at 37°C. The plates were washed and the horse radish peroxidase (HRP)-conjugated anti-human secondary antibody (anti-human Ig whole antibody from sheep (Amersham Life Science, NA933)) diluted 1:100 in dilution buffer was
10 added to the wells and incubated for 1 hour at 37°C. The plates were washed and OPD (o-phenylenediamine dihydrochloride) substrate with H₂O₂ in substrate buffer (50mM phosphate/25mM citrate, pH 7.2) was added to the wells. For a kinetics ELISA, the plate was read repeatedly (2 minute intervals for 15 minutes) unstopped (without "stop" buffer). Plates were read at 450nm.
15

Results

The results of the above experiment are presented in Table 9 and in Figure 12. The animals of group 2 received intranodal injections of ALVAC(2)-gp100(mod) followed by boosts with the modified gp100 peptides 209(2M) and 290(9V); the animals in group 3 received a subcutaneous injection of the
20 ALVAC(2) construct followed by peptide boosts; the animals in group 1 received intranodal injections of saline as a control.

As can be seen from Figure 12, both types of injection of the antigens induced a significant humoral response to the antigen.

In summary, the results of this Example demonstrate that injection of a
25 tumor antigen according to the invention induces both a significant humoral and cell mediated response.

EXAMPLE 5

This example presents data obtained from human melanoma patients primed with ALVAC(2)-gp100M and boosted with modified gp100 peptides
30 (g209-2M and g280-9V).

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Immunization Protocol

5 Patients were immunized subcutaneously in a prime-boost schedule with ALVAC(2)-gp100M ("prime"; lyophilized ALVAC(2)-gp100M resuspended in 1 ml of 0.4% NaCl; 0.5 ml injections (approximately $0.5 \times 10^{7.09}$ CCID₅₀ per injection)) and peptides g209-2M and g280-9V ("boost"; 1000 μ g/peptide in 1 ml total volume per week (0.2 ml/injection per day \times 5 days)). All patients: 1) were HLA-A0201 positive; 2) were between 18 and 70 years of age; 3) exhibited pathologically confirmed malignant melanoma; 4) demonstrated immunocompetence by reactivity to at least 2 or more out of 7 Cell Mediated Immunity (CMI) skin tests; 5) had blood hematology and chemistry values within the following ranges:

I) Hematology:

Hemoglobin	> 100g/L
Granulocytes	> 2.0 $\times 10^9$ /L
Lymphocytes	> 1.5 $\times 10^9$ /L
Platelets	> 100 $\times 10^9$ /L

II) Chemistry:

Serum creatinine	< 150 _mol/L
Serum total bilirubin	<30 _mol/L
AST, ALT, and ALP	Must be < 2x the normal upper limit or <5x the normal upper limit if due to liver metastases.

15

Patients "primed" with ALVAC(2)-gp100M on weeks 1, 4 and 7; "boosted" with peptides on weeks 10 and 13.

ELISPOT Analysis: These results are present in Tables 10 and 11. Peripheral Blood Mononuclear Cells ("PBMNC") were isolated by density centrifugation

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over Ficoll gradients. Cells were bulk-cultured at 3×10^6 /ml in AIM-V media along with a mixture of g209-2M and g280-9V or the HLA-A*0201 binding Flu peptide (all at 50 μ g/ml) for 8 days. IL-2 was added on days 3 and 5 of culture. On day 9, cells were harvested, counted and 2×10^5 cells/well plus 50 U/ml IL-2, 5 with and without the respective peptides, were plated in nitrocellulose membrane containing ELISPOT plates that had been precoated with anti-INF- γ antibodies. The plates were developed after 48 hours of culture. The numbers reported are the differences between the average of two wells restimulated with peptide and IL-2 and two wells treated only with IL-2.

10 Responses are the number of spots (counted by the electronic ELISPOT reader but confirmed in most cases by manual counting) per 2×10^5 PBMNC. The number of CD8+ T cells was not routinely determined but is typically 2-5-fold less than this number.

15 Having illustrated and described the principles of the invention in a preferred embodiment, it should be appreciated by those skilled in the art that the invention can be modified in arrangement and detail without departure from such principles. We claim all modifications coming within the scope of the following claims.

20 All publications, patents and patent applications referred to herein, are herein incorporated by reference in their entirety to the same extent as if each individual publication, patent or patent application was specifically and individually indicated to be incorporated by reference in its entirety.

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TABLE 1

Analysis of expression of gp100 antigen by ALVAC-gp100M

Human gp100M				
	Positive Plaques	negative plaques	total # of plaques	% positive
ALVAC std.	0	571	571	0
vCP1584	387	0	387	100
ALVAC gp100mod L	875	11	886	98.7

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TABLE 2

Group Number	Route of administration	Treatment days and compound administered	Number of Animals
1	Intranodal	Saline (NaCl 0.9%): days 28, 42, 56 Then 70, 71, 72, 73, 74 Then 84, 85, 86, 87 and 88	4
2	Intranodal	<u>ALVAC(2) - gp100 mod: days 28, 42, 56</u> *mgp100 peptides: days 70, 71, 72, 73, 74 Then 84, 85, 86, 87 and 88	4
3	Subcutaneous	Saline (NaCl 0.9%): day 1 <u>ALVAC(2) - gp100 mod: days 28, 42, 56</u> *mgp100 peptides: days 70 and 84	4

*209(2M)-IMDQVPFSY (SEQ.ID.NO.124); 290(9V) YLEPGPVTV (SEQ.ID.NO.125)

- 5 ▪ Group 1 animals (control) received the control article (saline for injection (NaCl 0.9%)).
- Group 3 animals received the control article (saline for injection (NaCl 0.9%)) on day 1 only.

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TABLE 3

Group Number	Dose level	Dose volume (ml/administration)
1	Saline (NaCl 0.9%): 0	0.250
2	Dose: 0.25×10^{74} CCID 50 ALVAC (2) - gp100 mod: 0.25×10^{74} CCID50 Dose: 200 µg (Total) of peptides IMDQVPFSY (209(2M)), and YLEPGPVTV (290(9V)) (100µg each)	0.250 0.2
3	Saline (NaCl 0.9%) ALVAC(2) - gp100 mod: 0.25×10^{74} CCID 50 Dose: 200 µg (Total) of peptides IMDQVPFSY (209(2M)), and YLEPGPVTV (290(9V)) (100µg each)	0.250 0.250 0.2

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TABLE 4

Days	Sites used
1 and 28	lower left
42	upper left
56	upper right
70	lower left
84	lower right

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TABLE 5

Peptide Pool #1

Peptide	Sequence	SEQ.ID.NO.
1329	HLAGVALLAVGATK	SEQ.ID.NO.3
1330	GALLAVGATKVPRNQ	SEQ.ID.NO.4
1331	VGATKVPRNQDWLGV	SEQ.ID.NO.5
1332	VPRNQDWLGVSQRQLR	SEQ.ID.NO.6
1333	DWLGVSRQLRTKAWN	SEQ.ID.NO.7
1334	SRLQRTKAWNRLQLYP	SEQ.ID.NO.8
1335	TKAWNRLQLYPEWTEA	SEQ.ID.NO.9
1336	RQLYPEWTEAQRLDC	SEQ.ID.NO.10
1337	EWTEAQRLDCWRGGQ	SEQ.ID.NO.11
1338	QRLLDCWRGGQVSLKV	SEQ.ID.NO.12
1339	WRGGQVSLKVSNNDGP	SEQ.ID.NO.13
1340	VSLKVSNDGFTLIGA	SEQ.ID.NO.14
1344	IALNFPGSQKVLPDG	SEQ.ID.NO.15
1345	PGSQKVLPDGQVIWV	SEQ.ID.NO.16
1346	VLPDGQVIWVNNTII	SEQ.ID.NO.17
1347	OVIWVNNTIINGSQV	SEQ.ID.NO.18
1348	NNTIINGSQVWGQQP	SEQ.ID.NO.19
1349	NGSQVWGQQPVYPQEQ	SEQ.ID.NO.20
1350	WGGQPVYPQETDDAC	SEQ.ID.NO.21
1351	VYPQETDDACIFPDG	SEQ.ID.NO.22
1352	TDDACIFPDGGPCPS	SEQ.ID.NO.23
1353	IFPDGGPCPSGSWSQ	SEQ.ID.NO.24
1355	GSWSQKRSFVVWKT	SEQ.ID.NO.25
1356	KRSFVYVWKTWGQYW	SEQ.ID.NO.26
1357	YVWKTWGQYWQVLGG	SEQ.ID.NO.27
1358	WGQYWQVLGGPVSGL	SEQ.ID.NO.28
1359	QVLGGPVSGLSIGTG	SEQ.ID.NO.29

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TABLE 6

Peptide Pool #2

Peptide	Sequence	SEQ.ID.NO.
1360	PVSGLSIGTGRAMLG	SEQ.ID.NO.30
1361	SIGTGRAMLCTHTM	SEQ.ID.NO.31
1362	RAMLGTHTMETVYH	SEQ.ID.NO.32
1363	THTMEVTYHRRGSR	SEQ.ID.NO.33
1364	VTVYHRRGSRSYVPL	SEQ.ID.NO.34
1365	RRGSRSYVPLAHSSS	SEQ.ID.NO.35
1366	SYVPLAHSSSAFTIT	SEQ.ID.NO.36
1368	AFTITDQVPFVSVS	SEQ.ID.NO.37
1369	DQVPFSVSQSQLRAL	SEQ.ID.NO.38
1370	SYSVSQQLRALDGGNK	SEQ.ID.NO.39
1372	DGGNKHFRLRNQPLTF	SEQ.ID.NO.40
1373	HFLRNQPLTFALQLH	SEQ.ID.NO.41
1374	QPLTFALQLHDPSGY	SEQ.ID.NO.42
1375	ALQLHDPSGYLAED	SEQ.ID.NO.43
1379	DFGDSSGTLISRALV	SEQ.ID.NO.44
1380	STGLISRALVVHTHY	SEQ.ID.NO.45
1381	SRALVVTHTYLEPGP	SEQ.ID.NO.46
1382	VVTHTYLEPGPVTAQV	SEQ.ID.NO.47
1383	LEPGPVTAQVVLQAA	SEQ.ID.NO.48
1384	VTAQVVVLQAAIPLTS	SEQ.ID.NO.49
1385	VLQAAIPLTCGSSP	SEQ.ID.NO.50
1386	IPLTSCGSSPVPGTT	SEQ.ID.NO.51
1388	VPGTTDGHRTAEP	SEQ.ID.NO.52
1389	DGHRTAEPNTTAG	SEQ.ID.NO.53
1390	TAEAPNTTAGQVPTT	SEQ.ID.NO.54
1392	QVPTTEVVGTTPQAA	SEQ.ID.NO.55
1393	EVVGTTPGQAPTAEP	SEQ.ID.NO.56

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TABLE 7

Peptide Pool #3

Peptide	Sequence	SEQ.ID.NO.
1394	TPGQAPTAEPSTTTS	SEQ.ID.NO.57
1395	PTAEPSTGTTSVQVPT	SEQ.ID.NO.58
1396	SGTTSVQVPTEVIS	SEQ.ID.NO.59
1397	VQVPTTEVISTAPVQ	SEQ.ID.NO.60
1398	TEVISTAPVQMPAAE	SEQ.ID.NO.61
1399	TAPVQMPAAESTGMMT	SEQ.ID.NO.62
1400	MPTAESTGMTPEKVP	SEQ.ID.NO.63
1401	STGMMTPEKVPVSEVM	SEQ.ID.NO.64
1402	PEKVPVSEVMGTTLA	SEQ.ID.NO.65
1403	VSEVMGTTLAEMSTP	SEQ.ID.NO.66
1404	GTTLAEMSTPEATGMI	SEQ.ID.NO.67
1405	EMSTPEATGMTPEAV	SEQ.ID.NO.68
1408	SIVVLSGTTAAQVTT	SEQ.ID.NO.69
1409	SGTTAAQVTTEWVE	SEQ.ID.NO.70
1410	AQVTTIEWVETTARE	SEQ.ID.NO.71
1411	TEWVETTAREPIPE	SEQ.ID.NO.72
1412	TTARELP:PEPEGPD	SEQ.ID.NO.73
1413	LPIPEPEGPDASSIM	SEQ.ID.NO.74
1414	PEGPDASSIMSTESI	SEQ.ID.NO.75
1415	ASSIMSTESITGSLG	SEQ.ID.NO.76
1416	STESITGSLGPLLDG	SEQ.ID.NO.77
1417	TGSLGPLLDGATLRL	SEQ.ID.NO.78
1418	PLLDGTATLRLVKRQ	SEQ.ID.NO.79
1419	TATLRLVKRQVPLDC	SEQ.ID.NO.80
1420	LVKRQVPLDCVLYRY	SEQ.ID.NO.81
1421	VPLDCVLYRYGSFSV	SEQ.ID.NO.82
1422	VLYRYGSFSVTLDIV	SEQ.ID.NO.83

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Table 8

Peptide Pool #4

Peptide	Sequence	SEQ.ID.NO.
1424	TLDIVQGIESAEILQ	SEQ.ID.NO.84
1425	QGIESAEILQAVPSG	SEQ.ID.NO.85
1426	AEILQAVPSGEGDAF	SEQ.ID.NO.86
1427	AVPSGEGDAFELTVS	SEQ.ID.NO.87
1428	EGDAFELTVSCQGL	SEQ.ID.NO.88
1429	ELTVSCQGLPKEAC	SEQ.ID.NO.89
1430	CQQGLPKEACMEIIS	SEQ.ID.NO.90
1431	PKEAACMEISSLPGCQP	SEQ.ID.NO.91
1432	MEISSLPGCOPPAQRL	SEQ.ID.NO.92
1434	PAQRLCQPVLPSPAC	SEQ.ID.NO.93
1435	CQPVLPSPACQLVLH	SEQ.ID.NO.94
1436	PSPACQLVLHQILKG	SEQ.ID.NO.95
1437	QLVLHQILKGSGTY	SEQ.ID.NO.96
1441	LADTNSLAVVSTQLI	SEQ.ID.NO.97
1442	SLAVVSTQLIMPQCQE	SEQ.ID.NO.98
1443	STQLIMPQCQEALGQ	SEQ.ID.NO.99
1444	MPGQEAGLGQVPLIV	SEQ.ID.NO.100
1445	AGLGQVPLIVGILLV	SEQ.ID.NO.101
1448	LMAVVVLASLIVRRRL	SEQ.ID.NO.102
1450	YRRRLMKQDFSVPQL	SEQ.ID.NO.103
1451	MKQDFSVPQLPHSSS	SEQ.ID.NO.104
1452	SVPQLPHSSSHWLRL	SEQ.ID.NO.105
1453	PHSSSHWLRLPRIFC	SEQ.ID.NO.106
1454	HWLRLPRIFCSCPIG	SEQ.ID.NO.107
1455	PRIFCSCPIGENSPL	SEQ.ID.NO.108

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TABLE 9

Monkey #	DAY (mOD/min)			
	0	57	68	96
1	3	5	2	2
2	4	6	12	10
3	7	6	10	8
4	7	6	8	8
5	5	9	20	15
6	11	8	10	12
7	11	23	51	30
8	7	30	70	22
9	1	7	5	3
10	2	6	6	4
11	3	7	14	8
12	6	9	15	6

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TABLE 10

Gp100-specific responses to g209-2M and g280-9V*

Patient	Pre 1 st Injection	Pre 2 nd Injection	Pre 3 rd Injection	Pre 4 th Injection	Pre 5 th Injection	4 wks post vaccination
#1	0	0	0	ND	ND	24.14
#2	0	14±2.8	54±6.4	16±7.8	ND	ND
#3	0	0	ND	ND	ND	ND
#4	0	0	24±13.4	1±2.1	ND	ND
#5	ND	6±6.4	ND	ND	ND	ND

5

TABLE 11

Flu-peptide specific responses*

Patient	Pre 1 st Injection	Pre 2 nd Injection	Pre 3 rd Injection	Pre 4 th Injection	Pre 5 th Injection	4 wks post vaccination
#1	>150	ND	>70	ND	ND	12.5
#2	ND	0	24	0	ND	ND
#3	23.5	7	ND	ND	ND	ND
#4	0	29	13.5	11.5	ND	ND
#5	ND	>200	ND	ND	ND	ND

* ND signifies that the values were not determined for the sample.

WE CLAIM:

1. An isolated and purified modified gp100 molecule capable of modulating an immune response in an animal.
5
2. A molecule according to claim 1 having a nucleic acid sequence shown in Figure 1 (SEQ.ID.NO.1).
3. A molecule according to claim 1 or 2 which comprises:
10 (a) a nucleic acid sequence as shown in Figure 1 (SEQ.ID.NO.1) wherein T can also be U;
 (b) nucleic acid sequences complementary to (a);
 (c) nucleic acid sequences which are homologous to (a) or (b);
 (d) a fragment of (a) to (c);
15 (e) a nucleic acid which will hybridize to (a) to (d) under stringent hybridization conditions; and
 (f) a nucleic acid molecule differing from any of the nucleic acids of (a) to (d) in codon sequences due to the degeneracy of the genetic code.
- 20 4. The nucleic acid of any one of claims 1-3 wherein the nucleic acid is selected from the group consisting of viral nucleic acid, plasmid, bacterial DNA, naked/free DNA, and RNA.
- 25 5. A viral nucleic acid of claim 4 wherein the virus is selected from adenovirus, alphavirus or poxvirus.
6. A poxvirus of claim 5 which is vaccinia, fowlpox, avipox, TROVAC, ALVAC, NYVAC or MVA.
- 30 7. The poxvirus of claim 6 which is ALVAC.

8. A composition comprising the nucleic acid of any one of claims 1-7 and a pharmaceutically acceptable diluent or carrier.
- 5 9. A composition according to claim 8 further comprising an adjuvant.
- 10 11. A cell comprising a nucleic acid according to any one of claims 1-7 wherein the cell expresses a polypeptide encoded by the nucleic acid.
12. A cell according to claim 10 wherein the cell is an antigen-presenting cell.
13. A recombinant virus comprising a virus into which is inserted a nucleic acid according to any one of claims 1-7 wherein the nucleic acid encodes for a polypeptide, the recombinant virus causing the expression of the polypeptide in an infected cell.
14. A recombinant virus into which is inserted a nucleic acid according to any one of claims 1-7 wherein the nucleic acid encodes for a polypeptide, wherein cells infected with the said recombinant virus are capable of eliciting an immune response directly against a member selected from the group consisting of:
 - (1) the polypeptide;
 - (2) a fragment of the polypeptide;
 - (3) a cell expressing the polypeptide or a fragment thereof; or
 - (4) cells binding the protein or fragment thereof.
15. A recombinant virus according to claim 13 or 14 selected from adenovirus, alphavirus, or poxvirus.

16. A recombinant virus according to claim 15 wherein the poxvirus is vaccinia, fowlpox, avipox, TROVAC, ALVAC, NYVAC or MVA.
- 5 17. A recombinant virus according to claim 16 wherein the virus is ALVAC.
18. A composition comprising a recombinant virus of any one of claims 13 to 17 and a pharmaceutically acceptable diluent or carrier.
- 10 19. An isolated protein encoded by a nucleic acid molecule according to any one of claims 1-7.
20. An isolated protein having the activity of a modified gp100 protein.
- 15 21. A protein having the amino acid sequence shown in Figure 2 (SEQ.ID.NO.2).
22. A method of modulating an animal's immune system comprising administering an effective amount of a gp100 or *gp100* which has been modified.
- 20 23. A method according to claim 22 where the gp100 is gp100M.
24. A method according to claim 22 wherein the *gp100* is *gp100M*.
- 25 25. A method according to claim 24 wherein the gp100M has a nucleic acid sequence as shown in Figure 1 (SEQ.ID.NO.1).
26. A method according to claim 23 wherein the gp100M has an amino acid shown in Figure 2 (SEQ.ID.NO.2).

27. A method of modulating an animal's immune system comprising administering to an animal in need thereof, an effective amount of a vector, into which has been inserted a *gp100* which has been modified, thereby modulating the animal's immune system.

5

28. A method according to claim 27 wherein the vector is administered with a lymphokine, cytokine, or a co-stimulatory molecule.

29. A method according to claim 28 wherein the cytokine is GM-CSF, IL-2,
10 IL-12, TNF, or IFN γ 1.

30. A method according to claim 28 wherein the molecule is a lymphokine.

31. A method according to claim 28 wherein the molecule is co-stimulatory
15 molecule.

32. A method according to claim 31 wherein the co-stimulatory molecule is a molecule of the B7 family.

20 33. A method according to any one of claims 27-32 wherein the vector is an adenovirus, alphavirus or poxvirus.

34. A method according to claim 33 wherein the poxvirus is vaccinia, fowlpox, avipox, TROVAC, ALVAC, NYVAC or MVA.

25

35. A method according to claim 34 wherein the poxvirus is ALVAC

36. A method for prophylactic treatment of cancer comprising administering to an animal an effective amount of a modified *gp100* or

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immunogenic fragment thereof, or a nucleic acid sequence encoding a modified gp100 or immunogenic fragment thereof.

37. A method according to claim 36 wherein the modified gp100 has an
5 amino acid sequence as shown in Figure 2 (SEQ.ID.NO.2).

38. A method according to claim 36 wherein the nucleic acid sequence is as
shown in Figure 1 (SEQ.ID.NO.1).

10 39. A method according to any one of claims 36, 37 or 38 wherein the cancer
is a melanoma.

40. A melanoma vaccine comprising a nucleic acid sequence encoding a
modified gp100.

15 41. A vaccine according to claim 40 wherein the modified gp100 is gp100M.

42. A vaccine according to claim 41 wherein the gp100M has the amino acid
sequence as shown in Figure 2 (SEQ.ID.NO.2).

20 43. A modified gp100 protein sequence which is modified to enhance its
binding to MHC molecules.

44. A modified protein sequence according to claim 43 wherein the protein
25 is gp100M.

45. The protein of claim 44 wherein the amino acid sequence is as shown in
Figure 2 (SEQ.ID.NO.2).

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46. A vaccine comprising a modified *gp100* nucleic acid sequence or its corresponding protein or protein fragment capable of eliciting the production of antibodies in a animal to corresponding antigens.
- 5 47. A vaccine according to claim 46 wherein the protein corresponding to the nucleic acid sequence is *gp100M*.
- 10 48. A vaccine according to claim 46 wherein the modified *gp100* nucleic acid sequence is *gp100M*.
- 15 49. A vaccine according to claim 48 wherein the *gp100M* nucleic acid sequence is as shown in Figure 1 (SEQ.ID.NO.1).
50. A vaccine according to claim 47 wherein the *gp100M* has an amino acid sequence as shown in Figure 2 (SEQ.ID.NO.2).
- 20 51. A vaccine comprising a modified *gp100* nucleic acid sequence or its corresponding protein or protein fragment capable of eliciting a cellular immune response.
52. A vaccine according to claim 51 wherein the protein corresponding to the nucleic acid sequence is *gp100M*.
- 25 53. A vaccine according to claim 51 wherein the modified *gp100* nucleic acid sequence is *gp100M*.
54. A vaccine according to claim 53 wherein the *gp100M* nucleic acid sequence is as shown in Figure 1 (SEQ.ID.NO.1).

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55. A vaccine according to claim 52 wherein the gp100M has an amino acid sequence is as shown in Figure 2 (SEQ.ID.NO.2).
56. An immunogenic composition containing a vaccine vector encoding for
5 a modified gp100 molecule.
57. A composition according to claim 56 wherein the modified gp100 molecule is gp100M.
- 10 58. A composition according to claim 57 wherein the modified gp100M has an amino acid sequence according to Figure 2 (SEQ.ID.NO.2).
59. A composition according to any one of claims 56, 57 or 58 wherein the vector is an adenovirus, alphavirus or poxvirus.
- 15 60. A composition according to claim 59 wherein the poxvirus is vaccinia, fowlpox, avipox, TROVAC, ALVAC, NYVAC or MVA.
61. A composition according to claim 60 wherein the poxvirus is ALVAC.
- 20 62. Immunogenic fragments of an isolated gp100M protein encoded by a nucleic acid molecule having a sequence according to SED ID NO. 1.

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FIGURE 1

ATGG ATCTGGTGCT AAAAGATGC CTTCTTCATT TGGCTGTGAT
 AGGTGCTTTC CTGGCTGTGG GGGCTACAAA AGTACCCAGA AACAGGACT GGCTTGGTGT
 CTCAGGCAA CTCAAGAACCA AAGCCTGGAA CAGGCAGCTG TATCCAGAGT GGACAGAAC
 CCAGAGACTT GACTGCTGGA GAGGTGGTCA AGTGTCCCTC AAGGTCACTA ATGATGGGCC
 TACACTGATT GGTCACCATG CCTCTCTTCTC TATTGCGGT AACTTCCCCTG GAAGCCAAAA
 GTTATTGCCA GATGGGCAGG TTATCTGGGT CAACAAATAC ATCATCAATG GGAGCAGGT
 GTGGGGAGGA CAGCCAGTGT ATCCCCAGGA AACTGACGAT GCCTGCATCT TCCCTGATGG
 TGGACCTTGC CCATCTGGCT CTGGTCTCA GAAGAGAAC TTGGTTATG TCTGGAAGAC
 CTGGGGCCAA TACTGGCAGG TTCTAGGGGG CCGCTGCTC GGGCTGACCA TTGGGACAGG
 CAGGCAATG CTGGGCACAC ACACGATGGA AGTAGCTGTC TACCATGCC GGGGATCCGG
 GAGCTATGTC CCTCTCTGCTC ATTCAGCTC AGCCCTCACCC ATTATGGACC AGGTGCCCTT
 CTCGGTGAGC GTGTCTTCACTG TGCGGGCCTT GGATGGAGGG AACRAAGCACT TCTCTGAGAAA
 TCAGCCTCTG ACCTTTGCCCTC TCCAGCTCCA TGACCCCCAGT GGCTATCTGG CTGAACTGTA
 CCTCTCTTAC ACCTGGGACT TTGAGACAG TAGTGGAACCT CTGATCTCTC GGGCACTTGT
 GGTCACTCAT ACTTACCTGG AGCTGGCCC AGTCACTGTT CAGGGTGGC TGCAGGCTGC
 CATTCCTCTC ACCTCTGTG GCTCTCCCC AGTCTCAGGC ACCACAGATG GGCACAGGCC
 AACTGCAAGG GCCCCCTAAAC CCACAGCTGG CCAAGTGCCT ACTACAGAAAG TTGTTGGTAC
 TACACCTGGT CAGGGGCCAA CTGCGAGGCC CTCAGGAAAC AGCATCTGTC AGGTGCCAAC
 CACTGAAGTC ATAAGCAGTC CACCTGTGCA GATGCCAACT GCAGAGAGCA CAGGTATGAC
 ACCTGAGAGG GTGGCAGTTT CAGGGTCACT GGGTACCCAA CTGGCAGAGA TGTCNAACTCC
 AGAGGCTACAA GGTTATGACAC CTGCGAGAGT ATCAATTGTC GTGCTTCTG TAACACAGC
 TGCAGGTTA ACAACTACAG AGTGGGTGGA GACCAACAGCT AGAGAGCTAC CTATCCCTGA
 GCCTGAAGGT CCAGATGCCA GCTCAATCAT GTCTACGGAA AGTATTACAG GTTCCCTGGG
 CCCCCCTGTC GATGGTACAG CCACCTTAAG GCTGGTGAAG AGACAACTGC CCCCCTGGATTG
 TGTCTGTAT CGATATGGTT CCTTTTCCCTG CACCCCTGGAC ATTGTCCAGG GTATTGAAAG
 TGCGGAGATC CTGCGAGGCTG TGCGCTCCCG TGAGGGGGAT SCATTGAGC TGACTGTGTC
 CTGCCAAAGG GGGTTGCCCA AGGAAGCTGCTC CATGGAGRTC TCATGCCAG GGTGCCAGCC
 CCCCTGCCAG CGGTGTCGCC AGCCTGTGCT ACCCAGGCCA GCTGCCAGC TGTTTCTGCA
 CCAGATACTG AAGGGTGGCT CGGGGACATA CTGCTCTCAAT GTGCTCTGG CTGATACCAA
 CAGCCCTGCCA GTGGTCAGCA CCACGCTTAT CATCCTGGT CAAGAGAGCAG GCTCTGGGCC
 GGTTCCGCTG ATCGTGGGCCA TCTTGTGGT GTGATGGCT GTGGTCTCTG CATCTCTGAT
 ATATAGGGCC AGACTATGCA AGCAAGACTT CTCCGTACCC CAGTGGCCAC ATAGCAGCAG
 TCACTGGCTG CGTCTACCCCCC GCATCTTCTG CTCTGTGCC ATTGGTGAAGA ACAGCCCCCT
 CCTCAGTGGG CAGCAGGTCT GA

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FIGURE 2

Met Asp Leu Val Leu Lys Arg Cys Leu Leu His Leu Ala Val Ile Gly		
1	5	10
Ala Leu Leu Ala Val Gly Ala Thr Lys Val Pro Arg Asn Gln Asp Trp		
20	25	30
Leu Gly Val Ser Arg Gln Leu Arg Thr Lys Ala Trp Asn Arg Gln Leu		
35	40	45
Tyr Pro Glu Trp Thr Glu Ala Gln Arg Leu Asp Cys Trp Arg Gly Gly		
50	55	60
Gln Val Ser Leu Lys Val Ser Asn Asp Gly Pro Thr Leu Ile Gly Ala		
65	70	75
Asn Ala Ser Phe Ser Ile Ala Leu Asn Phe Pro Gly Ser Gln Lys Val		
85	90	95
Leu Pro Asp Gly Gln Val Ile Trp Val Asn Asn Thr Ile Ile Asn Gly		
100	105	110
Ser Gln Val Trp Gly Gly Gln Pro Val Tyr Pro Gln Glu Thr Asp Asp		
115	120	125
Ala Cys Ile Phe Pro Asp Gly Gly Pro Cys Pro Ser Gly Ser Trp Ser		
130	135	140
Gln Lys Arg Ser Phe Val Tyr Val Trp Lys Thr Trp Gly Gln Tyr Trp		
145	150	155
Gln Val Leu Gly Gly Pro Val Ser Gly Leu Ser Ile Gly Thr Gly Arg		
165	170	175
Ala Met Leu Gly Thr His Thr Met Glu Val Thr Val Tyr His Arg Arg		
180	185	190
Gly Ser Arg Ser Tyr Val Pro Ile Ala His Ser Ser Ala Phe Thr		
195	200	205
Ile Met Asp Gln Val Pro Phe Ser Val Ser Val Ser Gln Ile Arg Ala		
210	215	220
Leu Asp Gly Gly Asn Lys His Phe Leu Arg Asn Gln Pro Leu Thr Phe		
225	230	235
Ala Leu Gln Leu His Asp Pro Ser Gly Tyr Leu Ala Glu Ala Asp Leu		
245	250	255
Ser Tyr Thr Trp Asp Phe Gly Asp Ser Ser Gly Thr Leu Ile Ser Arg		
260	265	270
Ala Leu Val Val Thr His Thr Tyr Leu Glu Pro Gly Pro Val Thr Val		
275	280	285
Gln Val Val Leu Gin Ala Ala Ile Pro Leu Thr Ser Cys Gly Ser Ser		
290	295	300
Pro Val Pro Gly Thr Thr Asp Gly His Arg Pro Thr Ala Glu Ala Pro		
305	310	315
Asn Thr Thr Ala Gly Gln Val Pro Thr Thr Glu Val Val Gly Thr Thr		
325	330	335
Pro Gly Gln Ala Pro Thr Ala Glu Pro Ser Gly Thr Thr Ser Val Gln		
340	345	350
Vai Pro Thr Thr Glu Val Ile Ser Thr Ala Pro Val Gln Met Pro Thr		
355	360	365

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FIGURE 2 (CONT'D)

Ala Glu Ser Thr Gly Met Thr Pro Glu Lys Val Pro Val Ser Glu Val
 370 375 380
 Met Gly Thr Thr Leu Ala Glu Met Ser Thr Pro Glu Ala Thr Gly Met
 385 390 395 400
 Thr Pro Ala Glu Val Ser Ile Val Val Leu Ser Gly Thr Thr Ala Ala
 405 410 415
 Gln Val Thr Thr Glu Trp Val Glu Thr Thr Ala Arg Glu Leu Pro
 420 425 430
 Ile Pro Glu Pro Glu Gly Pro Asp Ala Ser Ser Ile Met Ser Thr Glu
 435 440 445
 Ser Ile Thr Gly Ser Leu Gly Pro Leu Leu Asp Gly Thr Ala Thr Leu
 450 455 460
 Arg Leu Val Lys Arg Gln Val Pro Leu Asp Cys Val Leu Tyr Arg Tyr
 465 470 475 480
 Gly Ser Phe Ser Val Thr Leu Asp Ile Val Gln Gly Ile Glu Ser Ala
 485 490 495
 Glu Ile Leu Gln Ala Val Pro Ser Gly Glu Gly Asp Ala Phe Glu Leu
 500 505 510
 Thr Val Ser Cys Gln Gly Gly Leu Pro Lys Glu Ala Cys Met Glu Ile
 515 520 525
 Ser Ser Pro Gly Cys Gln Pro Pro Ala Gln Arg Leu Cys Gln Pro Val
 530 535 540
 Leu Pro Ser Pro Ala Cys Gln Leu Val Leu His Gln Ile Leu Lys Gly
 545 550 555 560
 Gly Ser Gly Thr Tyr Cys Leu Asn Val Ser Leu Ala Asp Thr Asn Ser
 565 570 575
 Leu Ala Val Val Ser Thr Gln Leu Ile Met Pro Gly Gln Glu Ala Gly
 580 585 590
 Leu Gly Gln Val Pro Leu Ile Val Gly Ile Leu Leu Val Leu Met Ala
 595 600 605
 Val Val Leu Ala Ser Leu Ile Tyr Arg Arg Arg Leu Met Lys Gln Asp
 610 615 620
 Phe Ser Val Pro Gln Leu Pro His Ser Ser Ser His Trp Leu Arg Leu
 625 630 635 640
 Pro Arg Ile Phe Cys Ser Cys Pro Ile Gly Glu Asn Ser Pro Leu Leu
 645 650 655
 Ser Gly Gln Gln Val
 660

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FIGURE 3

Nucleotide Sequence of C5H6gp100M

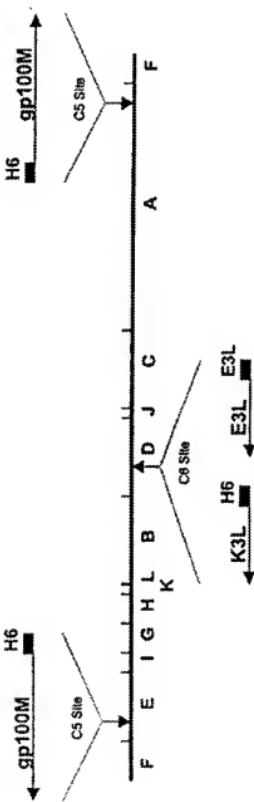
1-254 left C5 flanking arm
 255-376 H6 promoter
 377-2362 modified gp100 gene
 2363-2534 right C5 flanking arm

1 GGCTACTTTT CAACAAAGGA GCAGATGTA ACTACATCCT TGAAAGAAAT GGAAAATCAT
 61 ATACTGTGTTT GGAATTGATT AAAGGAAAGTT ACTCTGAGAC ACAAAAGAGG TAGCTGAAGT
 121 GGATCTCTA AAGGTACGGT ACTAAATTCAG TATAAAARGG ATCGTCGACG AGCTCGAATT
 181 CGGATCCGGG TTAAATTAATT AGTCATCAGG CAGGGCCGAGA ACAGAGACTAT CTGCTCGTTA
 241 ATTAATTTAGA GCTCTTCTTAT TCTATACTTA AAAAGTGTAA ATAATATCAA AGGTTCTTG
 301 GGGTTGTTGTT AAMTTGAAAG CGAGAAATAA TCATAAATTA ITTCATTATC GGATATCCG
 361 TTAAGTTGTT ATCGTATGG ATCTGGTGCt AAAAGAATGC CTTCTTCATT TGGCTGTGAT
 421 AGGTGCTTGT CTGGCTGTGG GGGCTACAAA AGTACCCCGA AACCAGGACT GGCTGGGTG
 481 CTCAGGCCA CTCAGAACCA AAGGCCGAA CAGGCCGCTG TATCCAGAGT GGACAGAAC
 541 CCAGAGACTT GACTGCTGGA GAGGTGGTCA AGTGTCCCCTC AAGGTCAAGTA ATGATGGCC
 601 TACATGATT GGTGCAATTT CCTCTCTCTC TATTGCTCTG AACTTCCCTG GAAGCCAAA
 661 GGTATTGCCA GATGGCAGG TTATCTGGGT CAACAAATTC ATCATCATTG GGAGCAAGT
 721 GTGGGGAGGA CAGCCAGTGT ATCCCCAGGA AACTGAGGCT GCCTGCTATCT TCCCTGTG
 781 TGGAGCTTGC CCATCTGGCT CTGGTCTCA GAAGAGAAGC TTGTTTATG TCTGGARAGC
 841 CTGGGGCCAA TACTGGCAAG TTCTAGGGGG CCCAGTCTC GGGCTGAGCA TTGGGACAGG
 901 CAGGGCAATG CTGGGCACAC ACACCGATGGA AGTGAATGTC TACCATGCGC GGGGATCCCG
 961 GAGCTATGTT CCTCTCTGCT ATTCCAGCTC AGCCCTTCACCC ATTATGGACCC AGGTGCTT
 1021 CTCCGGTGAGC GTGTCCTGGT TGCGGGCCCT TGATGGAGGG AACAGCACT TCTCTGAGAAA
 1081 TCAGCCTCTG ACCCTTGGCC TCCAGCTCCA TGACCCCAAGT GGCTATCTGG CTGAGGCTGA
 1141 CCTCTCTCTAC ACCCTGGGACT TTGAGAGACAG TAGTGGAAAC CTGATCTCTC GGGCACTTGT
 1201 GGTCACTCAT ACTTACCTGG AGCCCTGGCC AGTCACTGTT CAGGTGGTCC TGCAGGCTGC
 1261 CATTCCTCTC ACCTCTCTG GCTCTCTCCCC AGTCTCCAGG ACCACAGATG GGCAAGGCC
 1321 AACTCGACAG GCCCCCTAACCA CCACAGCTGG CCAAGTGCCT ACTACAGAAAG TTGTTGGTAC
 1381 TACACCTGGT CAGGGCCCAA CTGCGAGACCC CTCTGGAAAC ACATCTGTG AGGTGCCAC
 1441 CACTGAACCT ATAAGCCTG TGACCCAGTGC GATGCCAACT GCAGAGAGAC CAGGTATGAC
 1501 ACCTTGAGAG TGTCGAGCTTG CAGAGGTCTA GGGTACACCA CTGGCAGAGA TGTCAACTCC
 1561 AGAGGCTACAA GGTTAGACAC CTGCGAGAGT ATCAATTGTT GTGCTTCTG GAACCCACAGC
 1621 TGCAAGGTTA ACAACTACAG AGTGGGTGGA GACCAAGACCT AGAGAGCTAC CTATCCJIGA
 1681 GCTCTGGAGGT CCAGATGCCA GCTCAATCAT GTCTACGGAA AGTATTCAG GTTCTCTGG
 1741 CCCCTCTGCT GATGTCACAG CCACCTTAAG GCTGGTGAAG AGACAAGTCC CCCTGGATIG
 1801 TGTTCCTGTAT CGATATGTCG CTTTCTCCG GACCCCTGGGACT ATTGTCCAGG GTATGAAAG
 1861 TGCCGAGAGTC CTGCGAGCTG TGCCCTCCCG TGAGGGGGAT GCATTGAGC TGACTGTGTC
 1921 CTGCGAACGG GGGCTGCCA AGGAAGCTG CATGGAGATC TCATGCCAGG GGTGCCAGCC
 1981 CCCTGCCAGG CGGGCTGTGCG AGCCCTGTGCT ACCAGGCCCA GCCTGCCAGC TGTTCTGCA
 2041 CCAGATACTG AAGGGTGGCT CGGGGACATA CTGCTCTCAT GTGCTCTGG CTGATACCAA
 2101 CAGCGGCCA GTGGTCACCA CCCAGCTTAT CATGCTCTGGT CAAGAAGCAG GCCTGGGCCA
 2161 GTTCTGGCT ATCGTGGGCA TCTTGTGTT GTGATGAGT GTGGCTCTGGT CATCTCTGT
 2221 ATATAGGCGC AGACTATGAG AGCAAGACTT CTCCGTCACCC CAGTGGCCAC ATAGCAGCAG
 2281 TCACTGGCTG CGTCTACCCCT GCATCTCTG CTCTGTGCC ATTGGTGAGA ACAGCCCCCT
 2341 CCTCAGTGGG CAGCAGGCTG GATTTTATC TCGAGTCTG AATCGATCCC GGGTTTTTAT
 2401 GACTAGTTA TCACGGCCG TTATAAAAGAT CTAAATGCA TAATTCTAA ATAATGAAA
 2461 AAAAGTACAT CATGAGCRAA CGCTTAGTAT ATTTTACAT GGAGATTAAC GCTCTATACC
 2521 GTTCTATGTT TATT

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FIGURE 4

ALVAC(2)-gp100M (vCP1584)
(ALVAC Xhol Restriction Map)



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FIGURE 5

Oligonucleotide Primers

IDC5-1

CGT GCC ATG GCA CAC AAA AGA GGT AGC TGA A

IDC5-2

CCA GGC CGC CGC ACT AAC GCG TTG CTC ATG ATG

CSL

CAC AAA AGA GGT AGC TGA AGT

MEL 01

ATG GAT CTG GTG CTA AAA AGA

MEL 05

ACC TTG CCC ATC TGG CTC TTG

MEL 09

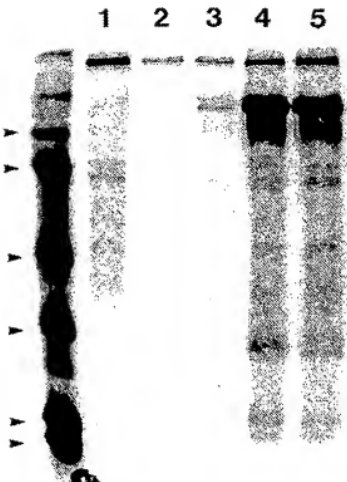
AGA TGC CAG CTC AAT CAT GTG

CSR

ATA GAT CTT TAT AAG CGG CCG

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FIGURE 6



Molecular Weight Markers: 200, 98.6, 68, 43, 29, 18, 14 kDa
Lane 1: Uninfected HeLa cells

Lane 2: HeLa cells infected with ALVAC

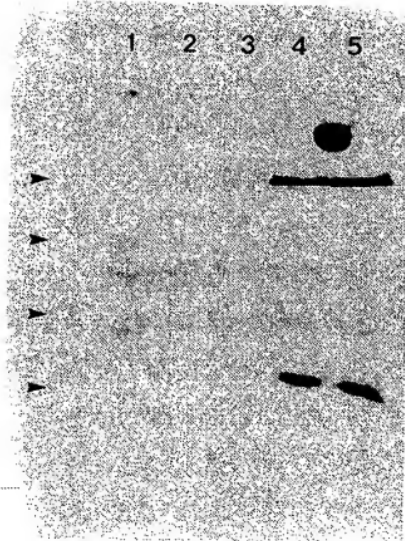
Lane 3: HeLa cells infected with ALVAC-gp100 (vCP1465)

Lane 4: HeLa cells infected with ALVAC(2)-gp100M (vCP1584)

Lane 5: HeLa cells infected with ALVAC(2)-gp100M (sister of vCP1584)

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FIGURE 7



Molecular Weight Markers: 97, 68, 43, 29 kDa

Lane 1: Uninfected HeLa cells

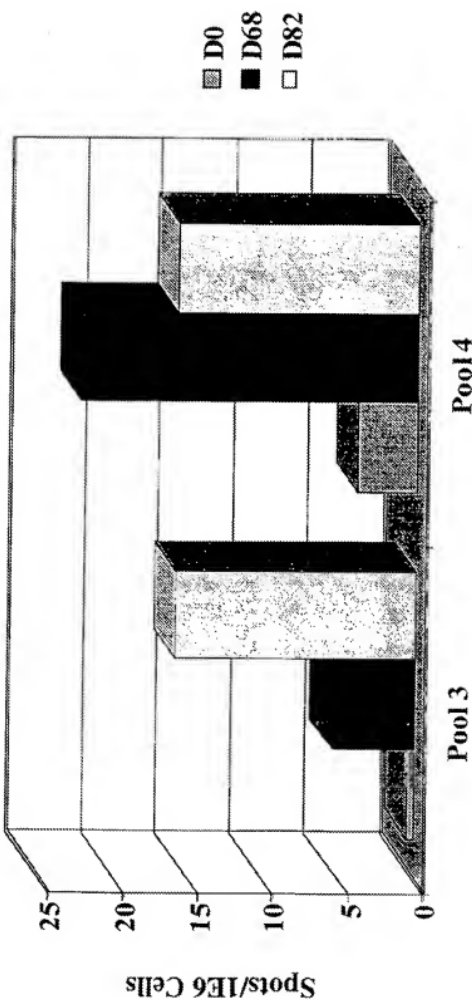
Lane 2: HeLa cells infected with ALVAC

Lane 3: HeLa cells infected with ALVAC-gp100 (vCP1465)

Lane 4: HeLa cells infected with ALVAC(2)-gp100M (vCP1584)

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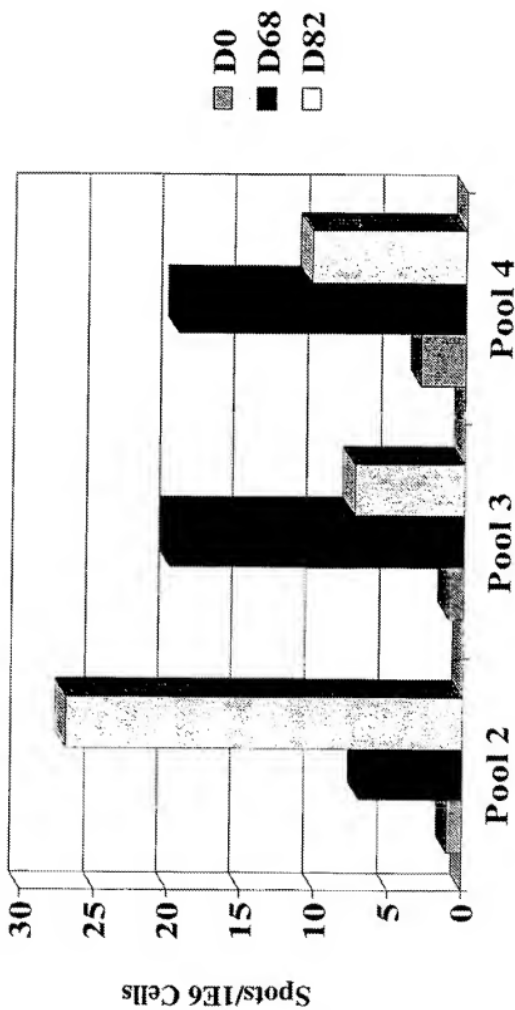
FIGURE 8
Monkey #6 (Intranodal Administration)



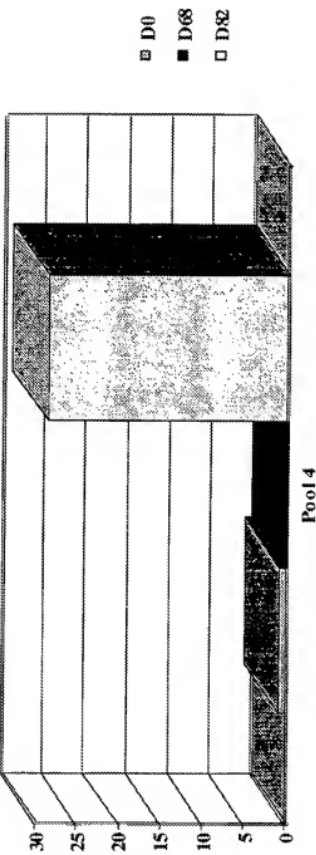
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FIGURE 9

Monkey #7 (Intranodal Administration)



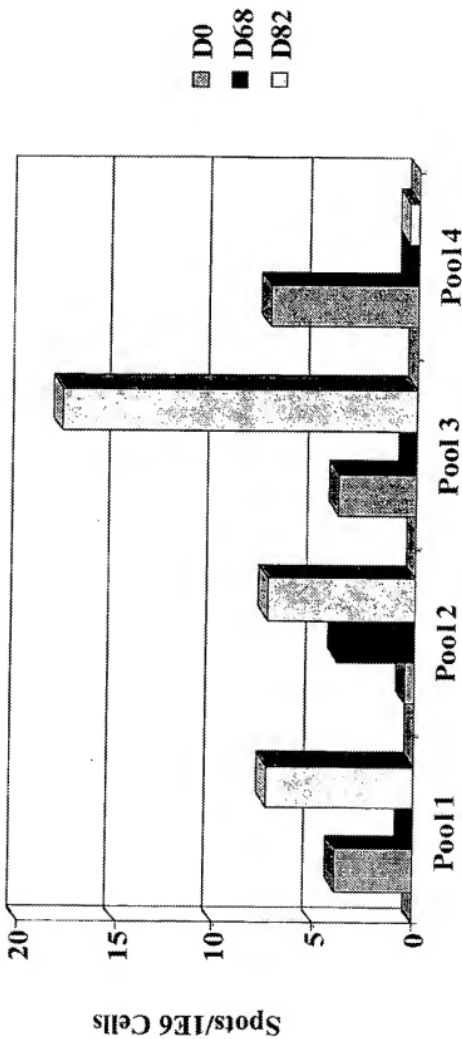
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FIGURE 10**Monkey # 11 (Subcutaneous Administration)**

Spots/1E6 Cells

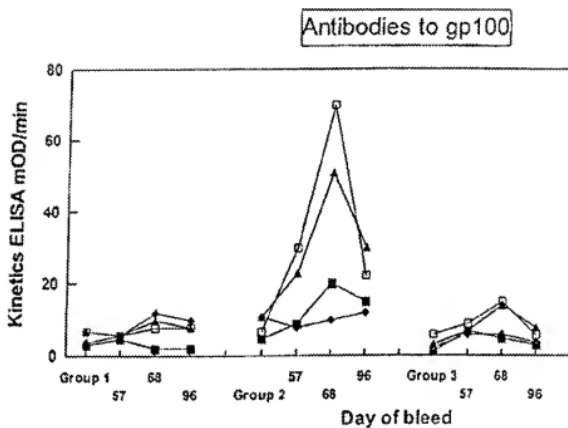
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FIGURE 11
Monkey #10 (Subcutaneous Administration)



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FIGURE 12



INTERNATIONAL SEARCH REPORT

International Application No
PCT/CA 00/01254

A. CLASSIFICATION OF SUBJECT MATTER

IPC 7 C07K14/705	C12N9/64	A61K39/00	C12N5/06	C12N7/00
A61K35/76	A61P35/00	A61K48/00		

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

MEDLINE, CANCERLIT, LIFESCIENCES, EMBASE, CHEM ABS Data, SCISEARCH, BIOSIS, WPI Data, EPO-Internal, PAJ

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>IRVINE KARI R ET AL: "Recombinant virus vaccination against "self" antigens using anchor-fixed immunogens." CANCER RESEARCH, vol. 59, no. 11, 1 June 1999 (1999-06-01), pages 2536-2540, XP002161590 ISSN: 0008-5472</p> <p>page 2536, left-hand column, line 39 -right-hand column, line 14 page 2356, right-hand column, line 47 -page 2357, left-hand column, line 1 table 1</p> <p>---</p> <p>-/-</p>	8,9, 13-16, 18-20, 22-24, 27-34, 36, 39-41, 43,44, 46-48, 51-53, 56,57, 59,60

 Further documents are listed in the continuation of box C. Patent family members are listed in annex.

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- *E* earlier document but published on or after the international filing date
- *L* document which may throw doubt on priority, claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- *O* document referring to an oral disclosure, use, exhibition or other means
- *P* document published prior to the international filing date but later than the priority date claimed

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INTERNATIONAL SEARCH REPORT

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